=>

Uploading C:\Program Files\Stnexp\Queries\323.str

L1 STRUCTURE UPLOADED

=> d

L1 HAS NO ANSWERS

L1

STR

G1 O, S, NH

Structure attributes must be viewed using STN Express query preparation.

=> s l1 full

REG1stRY INITIATED

Substance data SEARCH and crossover from CAS REGISTRY in progress... Use DISPLAY HITSTR (or FHITSTR) to directly view retrieved structures.

FULL SEARCH INITIATED 14:47:06 FILE 'REGISTRY'
FULL SCREEN SEARCH COMPLETED - 48452 TO ITERATE

100.0% PROCESSED 48452 ITERATIONS SEARCH TIME: 00.00.01

2066 SEA SSS FUL L1

2066 ANSWERS

SEARCH TIME: 00.00.01

=> s 13 and py<2001 20883874 PY<2001

280 L2

216 L3 AND PY<2001

=> s 14 and heterocy?

153465 HETEROCY?

L5 22 L4 AND HETEROCY?

=> d 1-22 ibib abs hitstr

L5 ANSWER 1 OF 22 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

2000:573791 CAPLUS

DOCUMENT NUMBER:

133:164009

TITLE:

L3

L4

Preparation of phenyl ureas and thioureas as orexin

receptor antagonists

INVENTOR(S):

Coulton, Steven; Johns, Amanda; Porter, Roderick Alan

PATENT ASSIGNEE(S):

Smithkline Beecham Plc, UK

SOURCE: PCT Int. Appl., 45 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

	PATENT NO.					KIND DATE		APPLICATION NO.										
	wo	2000	0475'	 77		A1 20000817		WO 2000-EP1150										
		W:	ΑE,	AL,	AM,	AT,	AU,	ΑZ,	BA,	BB,	BG,	BR,	BY,	CA,	CH,	CN,	CR,	CU,
			CZ,	DE,	DK,	DM,	EE,	ES,	FI,	GB,	GD,	GE,	GH,	GM,	HR,	HU,	ID,	IL,
			IN,	IS,	JP,	KE,	KG,	KP,	KR,	ΚZ,	LC,	LK,	LR,	LS,	LT,	LU,	LV,	MA,
			MD,	MG,	MK,	MN,	MW,	MX,	NO,	NZ,	PL,	PT,	RO,	.RU,	SD,	SE,	SG,	SI,
			SK,	SL,	TJ,	TM,	TR,	TT,	TZ,	UA,	ŪG,	US,	UZ,	VN,	YU,	ZA,	ZW,	AM,
									TJ,		_						•	
•		RW:	GH,	GM,	KE,	LS,	MW,	SD,	SL,	SZ,	TZ,	UG,	ZW,	AT,	BE,	CH,	CY,	DE,
			DK,	ES,	FI,	FR,	GB,	GR,	IE,	IT,	LU,	MC,	NL,	PT,	SE,	BF,	ВJ,	CF,
									ML,						-	-	-	
	EP 1150977															2	0000	210
		1150																
										GB,	GR,	IT,	LI,	LU,	NL,	SE,	MC,	PT,
						LV,			•	•	•	•		-	-		•	
	JP	2002							1029		JP 2	000-	5984	97		2	0000	210
		2745															0000	
	ES	2226	785			Т3		2005	0401		ES 2	000-	9063	24		2	0000	210
		6699									US 2						0020	
PRIOR	TTY	APP	LN.	INFO	. :						GB 1	999-	3266			A 1	9990	212
											GB 1	999-	2643	0		A 1	9991	108
											WO 2	000-	EP11	50	1	W 2	0000	210
OTHER	SC	URCE	(S):			MAR	PAT	133:	1640									
GI	. – –		, , .															

- * STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY AVAILABLE VIA OFFLINE PRINT *
- The title compds. [I; Z = O, S; R1 = alkyl, alkenyl, alkoxy, etc.; R2-R6 = alkyl, alkenyl, alkoxy, etc.; adjacent pair of R2-R6 together with the carbon atoms to which they are attached form (un)substituted carbocyclyl, heterocyclyl; R7 = alkyl, alkenyl, alkoxy, etc.; n = 0-3] and their pharmaceutically acceptable salts which are non-peptide antagonists of human orexin receptors, in particular orexin-1 receptors, were prepared E.g., treatment of 4-amino-2-methylquinoline with carbonyl diimidazole in CH2Cl2 followed by addition of 6-amino-2-methylbenzoxazole afforded II which showed pKb > 6.0 against orexin-1 receptor. In particular, compds. I are of potential use in the treatment of obesity including obesity observed in Type 2(non-insulin-dependent) diabetes patients and/or sleep disorders.

 IT 288150-79-8P
 - RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)
- (preparation of Ph ureas and thioureas as orexin receptor antagonists) RN 288150-79-8 CAPLUS
- CN 2-Propenamide, 3-[5-[[[(8-fluoro-2-methyl-4-quinolinyl)amino]carbonyl]amin o]-2-methoxyphenyl]-N-methyl- (9CI) (CA INDEX NAME)

IT 288151-91-7

RL: RCT (Reactant); RACT (Reactant or reagent) (preparation of Ph ureas and thioureas as orexin receptor antagonists)

RN 288151-91-7 CAPLUS

CN 2-Propenoic acid, 3-(2-methoxy-5-nitrophenyl)-, (2E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

IT 288151-85-9P 288151-86-0P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of Ph ureas and thioureas as orexin receptor antagonists)

RN 288151-85-9 CAPLUS

CN 2-Propenamide, 3-(2-methoxy-5-nitrophenyl)-N-methyl-, (2E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 288151-86-0 CAPLUS

2-Propenamide, 3-(5-amino-2-methoxyphenyl)-N-methyl-, (2E)- (9CI) (CA CN INDEX NAME)

Double bond geometry as shown.

REFERENCE COUNT: 7 THERE ARE 7 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 2 OF 22 CAPLUS COPYRIGHT 2006 ACS on STN

1999:551731 CAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER:

131:170173

TITLE:

Preparation of arylacrylate esters as precursors for

organoleptic compounds

INVENTOR(S): Anderson, Denise; Frater, Georg

PATENT ASSIGNEE(S): Givaudan Roure (International) S.A., Switz.

SOURCE:

Eur. Pat. Appl., 16 pp.

CODEN: EPXXDW

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATEN	NT NO.	KIND	DATE	APPLICATION NO.	DATE
	36211	A2	19990818	EP 1999-810036	19990119 <
		A3	19990825		
- F				GR, IT, LI, LU, NL	, SE, MC, PT,
	IE, SI, LT		•		
	38986	Α	20021130	IN 1999-MA51	
SG 93	3823	A 1	20030121	SG 1999-82	19990113
ZA 99	900567	Α	19990726	ZA 1999-567	
CN 12	227837	`A	19990908	CN 1999-101847	. 19990202 <
MX 99	901281	Α	20000731	MX 1999-1281	19990204 <
BR 99	900443	Α	20000502	BR 1999-443	19990210 <
AU 99	916430	A1	19991021	AU 1999-16430	
	25999		20001026		
	000063328	A2	20000229	JP 1999-33906	19990212 <
US 60	096918	Α	20000801	US 1999-249384	19990212 <
PRIORITY A	APPLN. INFO.:			EP 1998-810114	A 19980213
OTHER SOUP	RCE(S):	MARPAT	131:170173		
	• •			(aromatic) hydrocarl	bvl.
				bstituted by an ionic	
				rocarbyl, aryl, etc.	
				thylene; Z1 = CR2:CH	
				e (sic), (un)substitu	
				der use conditions to	
				ntimicrobial and/or of	
				Thus, 2-(HO)C6H4CHO	was condensed
with	Ph3P:CMeCO2Et	to give	(E)-2-(HO)C	6H4CH:CMeCO2Et.	
IT 23840	02-44-3P				

RL: MOA (Modifier or additive use); SPN (Synthetic preparation); PREP (Preparation); USES (Uses)

(preparation of arylacrylate esters as precursors for organoleptic compds.)

RN 238402-44-3 CAPLUS

CN 2-Butenoic acid, 3-[4-(diethylamino)-2-hydroxyphenyl]-, ethyl ester, (2E)-(9CI) (CA INDEX NAME)

Double bond geometry as shown.

L5 ANSWER 3 OF 22 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1998:479029 CAPLUS

DOCUMENT NUMBER:

129:122458

TITLE:

Preparation of N,N'-diphenylurea derivatives as

interleukin-8 receptor antagonists

INVENTOR(S):

Widdowson, Katherine Louisa; Veber, Daniel Frank; Jurewicz, Anthony Joseph; Hertzberg, Robert Philip;

Rutledge, Melvin Clarence, Jr.

PATENT ASSIGNEE(S):

Smithkline Beecham Corporation, USA

SOURCE:

U.S., 50 pp., Cont.-in-part of U.S. Ser. No. 641,990.

CODEN: USXXAM

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE	DATE		
					-		
us 5780483	Α	19980714	US 1996-701299	1996082	L <		
US 5886044	Α	19990323	US 1996-641990	1996032) <		
US 6211373	B1	20010403	US 1998-111663	1998070	3		
PRIORITY APPLN. INFO.:			US 1995-390260	B2 1995021	7.		
			US 1996-641990	A2 1996032)		
			WO 1996-US2260	W 1996021	5		
			US 1996-701299	A3 1996082	l		

OTHER SOURCE(S):

MARPAT 129:122458

GI

AB The title compds. [I; X = O, S; R = any functional moiety having an ionizable H and a pKa of ≤10 (sic); R1, Y = H, halo, NO2, cyano, (halo)alkyl, alkenyl, (halo)alkoxy, N3, HO, hydroxyalkyl, aryl, arylalkyl, aryloxy, arylalkoxy, heteroaryl, heteroarylalkyl, heterocyclylalkyl, heterocyclylalkoxy, arylalkenyl,

heteroarylalkenyl, (un) substituted NH2, CONH2, or SO3H, etc.; m, n = 1-3], which are useful for the treatment of disease states mediated by the chemokine, interleukin-8 (IL-8) (no data), are prepared Thus, Me 4-amino-3-hydroxybenzoate was added to a solution of Ph isocyanate in PhMe and the resulting mixture was stirred at .apprx.80° for 24-48 h to give 90% N-[2-hydroxy-4-(methoxycarbonyl)phenyl]-N'-phenylurea. 182499-23-6P 182499-25-8P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of N,N'-diphenylurea derivs. as interleukin-8 receptor antagonists for disease treatment)

RN 182499-23-6 CAPLUS

IT

CN 2-Propenoic acid, 3-[3-[[[(2-bromophenyl)amino]carbonyl]amino]-2-hydroxyphenyl]-, methyl ester, (2E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 182499-25-8 CAPLUS

CN 2-Propenamide, 3-[3-[[(2-bromophenyl)amino]carbonyl]amino]-2-hydroxyphenyl]-, (2E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

IT 86981-08-0P 182500-04-5P 182500-05-6P

182500-06-7P 182500-07-8P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of N,N'-diphenylurea derivs. as interleukin-8 receptor antagonists for disease treatment)

RN 86981-08-0 CAPLUS

CN 2-Propenoic acid, 3-(2-hydroxy-3-nitrophenyl)-, (2E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 182500-04-5 CAPLUS

CN 2-Propenoic acid, 3-(2-hydroxy-3-nitrophenyl)-, methyl ester, (2E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 182500-05-6 CAPLUS

CN 2-Propenoic acid, 3-(3-amino-2-hydroxyphenyl)-, methyl ester, (2E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 182500-06-7 CAPLUS

CN 2-Propenamide, 3-(2-hydroxy-3-nitrophenyl)-, (2E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 182500-07-8 CAPLUS

CN 2-Propenamide, 3-(3-amino-2-hydroxyphenyl)-, (2E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

REFERENCE COUNT:

84 THERE ARE 84 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

CAPLUS COPYRIGHT 2006 ACS on STN ANSWER 4 OF 22

ACCESSION NUMBER:

1997:679050 CAPLUS

DOCUMENT NUMBER:

127:346406

TITLE:

Preparation of acylaminocinnamates and related

compounds as integrin antagonists.

INVENTOR(S):

Chen, Barbara B.; Chen, Helen Y.; Clare, Michael;

Docter, Stephen H.; Khanna, Ish Kumar; Koszyk, Francis

Jan; Malecha, James W.; Miyashiro, Julie M.; et al.

PATENT ASSIGNEE(S):

SOURCE:

G.D. Searle and Co., USA PCT Int. Appl., 278 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PA							KIND DATE			APPLICATION NO.					DATE		
WO	9736	860					1997	1009	WO 1997-US4462					19	9970:	325 <	
	W:	AL,	AM,	AT,	AU,	ΑZ,	BA,	BB,	BG,	BR,	BY,	CA,	CH,	CN,	CU,	CZ,	DE,
							GE,										
							LU,										
							SG,										
							KG,						,	,		•	•
	RW:		•	•	•		SZ,	•	•		•		DK.	ĖS.	FI.	FR.	GB.
							NL,										
	•	•	•	•	SN,	•	•	,	,	,	. 20,	,	00,	,	 ,	J,	 ,
CA	CA 2250690					•		1009		CA 1	997-	2250:	690		1 '	9970:	325 <
	8940						1999										325 <
	8940										,	J _ U					,
									GB	GR	тт	T.T	T.JT	NT.	SE	РΨ	IE, FI
JT.	2000							-						•			325 <
	2197				E		2002										
	2179	-															
							1997										
	AU 9723371 PRIORITY APPLN. INFO.:						1997	1022			997- 996-						326 < - -
PRIORIT	I APP	TM.	INFO	. :													
OM11777 ~							107	2464		WO I	997-	US44	02	1	M T;	9970.	323
OTHER SOURCE(S):				MAR.	PAT	12/:	3464	Ub				•					

GI.

Title compds. [I; A = NR5C(Y1)NR7R8, NR5C(NR7)Y2; Y1 = NR2, O, S; R = XR3; AΒ R1 = H, alkyl, amino, acylamino, etc.; X = O, S, NR4; R2 = H, (substituted) alkyl, aryl, OH, alkoxy, cyano, NO2, amino, aminocarbonyl, alkenyl, alkynyl, etc.; R3, R4 = H, alkyl, alkenyl, alkynyl, haloalkyl, aryl, aralkyl, sugar residue, steroid residue, etc.; R5 = H, alkyl, alkenyl, alkynyl, PhCH2, PhCH2CH2; R7 = H, (substituted) alkyl, alkenyl, alkynyl, aralkyl, cycloalkyl, bicycloalkyl, aryl, acyl, etc.; R50 = H, alkyl, (substituted) aryl, etc.; R52 = H, acylamino, (substituted) hydrazino; R2R7 = (substituted) heterocyclyl, heteroaryl; R7R8 = (substituted) heterocyclyl; Y2R7 = (substituted) heterocyclyl; Z1, Z2, Z3, Z5 = H, alkyl, OH, alkoxy, aryloxy, aralkoxy, halo, haloalkyl, haloalkoxy, NO2, amino, aminoalkyl, cyano, alkylsulfonyl, carboxyalkenyl, (fused) aryl, etc.; B = (CH2)pO, CH:CH, CH2CONH, CONH(CH2)p, CO2, SO2NH, etc.; m = 0-2; n = 0-3; p = 0-2]. Thus, 3-[2-methoxy-4-[[[3-[(1,2,3,4-tetrahydropyrimidin-2yl)amino]phenyl]carbonyl]amino]phenyl]propionic acid trifluoroacetate (preparation given) antagonized $\alpha v\beta 3$ with IC50 = 0.43 nM. 198193-15-6P 198193-16-7P 198193-18-9P IT 198193-19-0P 198193-54-3P 198193-55-4P 198193-62-3P 198193-63-4P 198193-72-5P 198193-73-6P RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use);

Ι

BIOL (Biological study); PREP (Preparation); USES (Uses) (preparation of acylaminocinnamates and related compds. as integrin antagonists)

RN 198193-15-6 CAPLUS

CN 2-Propenoic acid, 3-[4-[[3-[(aminoiminomethyl)amino]benzoyl]amino]-2-methoxyphenyl]-, ethyl ester, (E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 198193-16-7 CAPLUS

CN 2-Propenoic acid, 3-[4-[[3-[(aminoiminomethyl)amino]benzoyl]amino]-2-methoxyphenyl]-, ethyl ester, (E)-, trifluoroacetate (10:11) (9CI) (CA INDEX NAME)

CM 1

CRN 198193-15-6 CMF C20 H22 N4 O4

Double bond geometry as shown.

CM 2

CRN 76-05-1 CMF C2 H F3 O2

RN 198193-18-9 CAPLUS

CN 2-Propenoic acid, 3-[4-[[3-[(aminoiminomethyl)amino]benzoyl]amino]-2-methoxyphenyl]-, (2E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 198193-19-0 CAPLUS

CN 2-Propenoic acid, 3-[4-[[3-[(aminoiminomethyl)amino]benzoyl]amino]-2-methoxyphenyl]-, (E)-, trifluoroacetate (5:6) (9CI) (CA INDEX NAME)

CM 1

CRN 198193-18-9 CMF C18 H18 N4 O4

Double bond geometry as shown.

CM 2

CRN 76-05-1 CMF C2 H F3 O2

RN 198193-54-3 CAPLUS

CN 2-Propenoic acid, 3-[2-[(ethoxycarbonyl)amino]-4-[[3-[(1,4,5,6-tetrahydro-2-pyrimidinyl)amino]benzoyl]amino]phenyl]-, (E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 198193-55-4 CAPLUS

CN 2-Propenoic acid, 3-[2-[(ethoxycarbonyl)amino]-4-[[3-[(1,4,5,6-tetrahydro-2-pyrimidinyl)amino]benzoyl]amino]phenyl]-, (E)-, trifluoroacetate (5:9) (9CI) (CA INDEX NAME)

CM 1

CRN 198193-54-3 CMF C23 H25 N5 O5

Double bond geometry as shown.

CM 2

CRN 76-05-1 CMF C2 H F3 O2

RN 198193-62-3 CAPLUS

CN 2-Propenoic acid, 3-[2-methoxy-4-[[3-[(1,4,5,6-tetrahydro-2-pyrimidinyl)amino]benzoyl]amino]phenyl]-, (2E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 198193-63-4 CAPLUS

CN 2-Propenoic acid, 3-[2-methoxy-4-[[3-[(1,4,5,6-tetrahydro-2-pyrimidinyl)amino]benzoyl]amino]phenyl]-, (E)-, trifluoroacetate (5:6) (9CI) (CA INDEX NAME)

CRN 198193-62-3 CMF C21 H22 N4 O4

Double bond geometry as shown.

CM 2

CRN 76-05-1 CMF C2 H F3 O2

RN 198193-72-5 CAPLUS

CN 2-Propenoic acid, 3-[4-[[3-[(aminoiminomethyl)amino]-5-(trifluoromethyl)benzoyl]amino]-2-methoxyphenyl]-, (E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 198193-73-6 CAPLUS

CN 2-Propenoic acid, 3-[4-[[3-[(aminoiminomethyl)amino]-5-(trifluoromethyl)benzoyl]amino]-2-methoxyphenyl]-, (E)-, trifluoroacetate

(2:3) (9CI) (CA INDEX NAME)

CM 1

CRN 198193-72-5

CMF C19 H17 F3 N4 O4

Double bond geometry as shown.

CM 2

CRN 76-05-1 CMF C2 H F3 O2

IT 198194-94-4P 198194-95-5P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of acylaminocinnamates and related compds. as integrin

antagonists)

RN 198194-94-4 CAPLUS

CN 2-Propenoic acid, 3-[2-[(ethoxycarbonyl)amino]-4-nitrophenyl]-,

1,1-dimethylethyl ester, (E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

198194-95-5 CAPLUS RN

2-Propenoic acid, 3-[2-[(ethoxycarbonyl)amino]-4-[[3-[(1,4,5,6-tetrahydro-CN 2-pyrimidinyl)amino]benzoyl]amino]phenyl]-, 1,1-dimethylethyl ester, (E)-(9CI) (CA INDEX NAME)

Double bond geometry as shown.

CAPLUS COPYRIGHT 2006 ACS on STN ANSWER 5 OF 22

ACCESSION NUMBER:

1997:41948 CAPLUS

DOCUMENT NUMBER:

126:59875

TITLE:

Preparation of beta-heterocyclyl-alpha,

beta-unsaturated ketone derivatives as inhibitors of

interleukin 1 production

INVENTOR(S):

Tanaka, Masayuki; Okita, Makoto; Miyamoto, Mitsuaki; Kaneko, Toshihiko; Kawahara, Tetsuya; Akamatsu, Keishi; Chiba, Kenichi; Obaishi, Hiroshi; Sakurai, Hideki; Abe, Shinya; Kobayashi, Seiichi; Yamanaka,

JP 1995-142394

A 19950518

Takashi

PATENT ASSIGNEE(S):

SOURCE:

Eisai Co., Ltd., Japan PCT Int. Appl., 254 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

LANGUAGE:

Patent Japanese

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
				-
WO 9636608	A1	19961121	WO 1996-JP1330	19960520 <

W: CA, US

RW: AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE 19961126 JP 1995-142394 19950518 <--JP 08311032 A2

PRIORITY APPLN. INFO .:

MARPAT 126:59875

OTHER SOURCE(S): For diagram(s), see printed CA Issue. GΙ

 α , β -Unsatd. ketone derivs. represented by general formula AB RCH: CHCOR1 [R = Q, Q1; wherein Z = NH, O, S; ring B = an optionally substituted aromatic ring; R2 = H, halo, optionally halogenated lower alkyl, etc.; R3 = H, optionally halogenated lower alkyl, cycloalkyl optionally having heteroatom(s), alkoxyalkyl, optionally substituted aryl, optionally substituted heteroaryl, etc.; R1 = CR4R5R6; wherein R4, R5 = H, optionally halogenated lower alkyl, etc.; R6 = H, optionally halogenated lower alkyl, cycloalkyl optionally having heteroatom(s), optionally substituted aryl, optionally substituted heteroaryl, etc.] or pharmacol. acceptable salts

thereof, which are useful for the prevention and treatment of interleukin 1 production-related diseases, e.g. inflammation, are prepared Thus, 1.68 g 7-ethyl-4-methoxymethoxy-3,5,8-trimethoxy-2-quinolinecarboxaldehyde and 1.0 g 3-hydroxy-3-methyl-2-butanone were dissolved in MeOH, treated with 0.21 g LiOH.H2O and heated at 50-60° for 1 h to give, after treatment of the product with 1 N aqueous HCl in EtOAc, the title quinolinylbutenone derivative (I; R7 = R1O = OMe, R8 = H, R9 = Et, R11 = CMe2OH). The latter compound and I (R7 = R9 = R1O = H, R8 = C1, R2 = R11 = Me) showed IC50 of 1.08 and <0.1 nM, resp., for inhibiting the production of interleukin 1α in human peripheral monocyte and 0.92 and <0.1 nM, resp., for inhibiting the production of interleukin 1β in human peripheral monocyte.

IT 185207-34-5P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of $\beta\text{--}$ heterocyclyl- $\alpha\text{,}$ $\beta\text{--}unsatd.$ ketone derivs. as inhibitors of interleukin 1 production)

RN 185207-34-5 CAPLUS

CN 2-Propenoic acid, 3-(2,5-dimethoxy-3-nitrophenyl)-, ethyl ester (9CI) (CA INDEX NAME)

L5 ANSWER 6 OF 22 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1996:643902 CAPLUS

DOCUMENT NUMBER:

125:275430

TITLE:

Preparation of N,N'-diphenylurea derivatives as

interleukin-8 receptor antagonists

INVENTOR(S):

Widdowson, Katherine Louisa; Veber, Daniel Frank; Jurewicz, Anthony Joseph; Rutledge, Melvin Clarence,

Jr.; Hertzberg, Robert Philip

PATENT ASSIGNEE(S):

Smithkline Beecham Corporation, USA

SOURCE:

PCT Int. Appl., 116 pp. CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

: 5

PATENT INFORMATION:

PATENT NO.	KIND DATE	APPLICATION NO.	DATE		
WO 9625157 W: JP, US	A1 19960822	WO 1996-US2260			
RW: AT, BE, C		GB, GR, IE, IT, LU, MC, EP 1996-906547			
JP 11503110		JP 1996-525199			
WO 9729743 ·	Al 19970821	CA 1996-2432662 WO 1996-US13632	19960821 <		
KP, KR, L	K, LR, LT, LV, MD,	CN, CZ, EE, GE, HU, IL, MG, MK, MN, MX, NO, NZ, VN, AM, AZ, BY, KG, KZ,	PL, RO, SG,		

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RW: KE, LS, MW, SD, SZ, UG, AT, BE, CH, DE, DK, ES, FI, FR, GB, GR,
             IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, ML,
             MR, NE, SN, TD, TG
                                19970902
                                             AU 1996-69007
                                                                     19960821 <--
    AU 9669007
                          A1
    AU 725456
                          B2
                                20001012
                                                                     19960821 <--
    EP 896531
                          A1
                                 19990217
                                             EP 1996-929723
        R: AT, ES, GR, LU, SE, MC, PT, IE, SI, LT, LV, FI
    CN 1215990
                                                                     19960821 <--
                          Α
                                 19990505
                                             CN 1996-180245
                                                                     19960821 <--
    JP 2000504722
                                             JP 1997-529318
                          T2
                                 20000418
    NZ 316710
                                                                   . 19960821 <--
                                             NZ 1996-316710
                                 20000526
                          Α
                                                                     19960821 <--
                                             BR 1996-12779
    BR 9612779
                          Α
                                 20001024
                                                                     19960821
                                             CN 2004-10032423
    CN 1539816
                          Α
                                20041027
                                             US 1997-894291
                                                                     19970815 <--
    US 6005008
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                                 19991221
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    US 6211373
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                                             US 1998-111663
                                 19981014
                                             NO 1998-3737
                                                                     19980814 <--
    NO 9803737
                          Α
    US 6180675
                          В1
                                 20010130
                                             US 1999-240354
                                                                     19990129
                                             US 1995-390260
                                                                  A2 19950217
PRIORITY APPLN. INFO .:
                                             WO 1996-US2260
                                                                     19960216
                                                                  W
                                                                  A3 19960320
                                             US 1996-641990
                                             CA 1996-2245927
                                                                  A3 19960821
                                             US 1996-701299
                                                                  A3 19960821
                                             WO 1996-US13632
                                                                  W
                                                                     19960821
```

OTHER SOURCE(S):

MARPAT 125:275430

I

GΙ

The title compds. [I; X = 0, S; R =any functional moiety having an ·AB ionizable H and a pKa of ≤10; R1, Y = H, halo, NO2, cyano, C1-10 (halo)alkyl, C2-10 alkenyl, C1-10 (halo)alkoxy, N3, H0, C1-4 hydroxyalkyl, aryl, aryl-C1-4 alkyl, aryloxy, aryl-C1-4 alkoxy, heteroaryl, heteroarylalkyl, heterocyclyl, heterocyclyl-C1-4 alkyl, heterocyclyl-C1-4 alkoxy, aryl-C2-10 alkenyl, heteroaryl-C2-10 alkenyl, (un) substituted NH2, carbamoyl, or SO3H, etc.; m, n = 1-3], which are useful for the treatment of disease states mediated by the chemokine, interleukin-8 (IL-8) (no data), are prepared The chemokine-mediated disease is selected from psoriasis or atopic dermatitis, asthma, chronic obstructive pulmonary disease, adult respiratory distress syndrome, arthritis, inflammatory bowel disease, Crohn's disease, ulcerative colitis, septic shock, endotoxic shock, gram neg. sepsis, toxic shock syndrome, stroke, cardiac and renal reperfusion injury, glomerulo-nephritis, thrombosis, Alzheimer's disease, graft vs. host reaction, and allograft rejections. Thus, 1.19 mmol Me 4-amino-3-hydroxybenzoate was added to a solution of 1.19 mmol Ph isocyanate in toluene and the resulting mixture was stirred at .apprx.80° for 24-48 h to give 90% N-[2-hydroxy-4-(methoxycarbonyl)phenyl]-N'-phenylurea. 182499-23-6P 182499-25-8P IT

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of N,N'-diphenylurea derivs. as interleukin-8 receptor antagonists for disease treatment)

RN 182499-23-6 CAPLUS

CN 2-Propenoic acid, 3-[3-[[[(2-bromophenyl)amino]carbonyl]amino]-2-hydroxyphenyl]-, methyl ester, (2E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 182499-25-8 CAPLUS

CN 2-Propenamide, 3-[3-[[[(2-bromophenyl)amino]carbonyl]amino]-2-hydroxyphenyl]-, (2E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

IT 86981-08-0P 182500-04-5P 182500-05-6P

182500-06-7P 182500-07-8P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of N,N'-diphenylurea derivs. as interleukin-8 receptor antagonists for disease treatment)

RN 86981-08-0 CAPLUS

CN 2-Propenoic acid, 3-(2-hydroxy-3-nitrophenyl)-, (2E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 182500-04-5 CAPLUS

CN 2-Propenoic acid, 3-(2-hydroxy-3-nitrophenyl)-, methyl ester, (2E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 182500-05-6 CAPLUS

CN 2-Propenoic acid, 3-(3-amino-2-hydroxyphenyl)-, methyl ester, (2E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 182500-06-7 CAPLUS

CN 2-Propenamide, 3-(2-hydroxy-3-nitrophenyl)-, (2E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 182500-07-8 CAPLUS

CN 2-Propenamide, 3-(3-amino-2-hydroxyphenyl)-, (2E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

L5 ANSWER 7 OF 22 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1993:244465 CAPLUS

DOCUMENT NUMBER: 118:244465

TITLE: Silver halide photographic light-sensitive material

INVENTOR(S): Matushita, Tetunori

PATENT ASSIGNEE(S): Fuji Photo Film Co., Ltd., Japan

SOURCE: Eur. Pat. Appl., 74 pp.

CODEN: EPXXDW

DOCUMENT TYPE: Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
EP 508432	A1	19921014	EP 1992-106180	19920409 <
EP 508432	B1	19980325		
R: DE, FR, GB,	NL			
JP 04311952	A2	19921104	JP 1991-103584	19910410 <
US 5266453	Α	19931130	US 1992-866517	19920410 <
PRIORITY APPLN. INFO.:			JP 1991-103584 A	19910410
OTHER SOURCE(S):	MARPAT	118:244465		
GI				

$$\begin{bmatrix} (R^{1})_{n_{1}} - Y^{1} \end{bmatrix}_{n_{0}} \qquad \qquad (Z)_{h}$$

$$R^{2} \qquad \qquad X - D - M \qquad (Z)$$

AB Photog. material with improved safelight property contains in ≥1 hydrophilic colloidal layer ≥1 filter dye which is irreversibly bleached during processing step. The filter dye comprises I (R1,R2 = H, or a substitutable) group; n0, n1, n2 = 0-1; h = 1-2; R1,R2,R3 = may together form a hydrocarbon or heterocyclic ring; Y1 = CO, CO(NR4), CS, C(N+R5R6), SO, SO2, C(CR7R8), R6CN, or C6CCR9 in [(R1)n1 Y1] when n1 = 1 and in Y1(R3)n2 when n2 = 1 in which R4-R9 = H or a substitutable group, Y1 = CN, NO2 in [(R1)nY1] when n1 = 0 and in Y1(R3)n2 when n2 = 0; x - divalent linkage; D = photog. dye residue; M = amphoteric group.

IT 146844-68-0

RL: USES (Uses)

(photog. material with improved safelight property containing filter dye of)

RN 146844-68-0 CAPLUS

1-Butanaminium, N-[2-[[4-[3-[[4-[[5-chloro-1,2,3,6-tetrahydro-3-methyl-1-[2-(octyloxy)-2-oxoethyl]-2,6-dioxo-4-pyrimidinyl]oxy]phenyl]amino]-2-cyano-3-oxo-1-propenyl]-3-methoxyphenyl]ethylamino]ethyl]-N,N-dimethyl-4-sulfo-, inner salt (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} & \text{Me} \\ & + \\ -\text{CH}_2 - \text{CH}_2 - \text{N}^+ & (\text{CH}_2) \text{ 4} - \text{SO}_3^- \\ & & \\ & \text{Me} \end{array}$$

CAPLUS COPYRIGHT 2006 ACS on STN ANSWER 8 OF 22

ACCESSION NUMBER:

1993:29821 CAPLUS

DOCUMENT NUMBER:

118:29821

TITLE:

Photographic material containing quick bleachable

dyes

INVENTOR(S):

Kawashima, Yasuhiko; Yamauchi, Reiko; Kagawa, Nobuaki

PATENT ASSIGNEE(S):

SOURCE:

Konica Co., Japan Jpn. Kokai Tokkyo Koho, 37 pp.

CODEN: JKXXAF

DOCUMENT TYPE:

Patent

LANGUAGE:

Japanese

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE		
JP 04116639 PRIORITY APPLN. INFO.: GI	A2	19920417	JP 1990-237765 JP 1990-237765	19900907 < 19900907		

Ι

$$R^{17}$$
 $C = CH - (CH = CH)_{m}$
 R^{13}
 R^{11}
 R^{12}
 R^{15}

$$c = cH - cH = cH$$
 $n = cH$
 $n = cH$

$$C = CH - (CH = CH) \xrightarrow{R^{33}} \xrightarrow{R^{31}} \xrightarrow{R^{32}} \xrightarrow{R^{34}} \xrightarrow{R^{35}} \xrightarrow{R^{35}} \xrightarrow{R^{31}} \xrightarrow{R^{35}} \xrightarrow{R^{$$

The title photog. material contains a dispersed fine solid powder of a AB compound selected from I, II and III [R1,2 = H, (cyclo)alkyl, alkenyl, aryl, heterocyclyl, acyl, sulfonyl; R1 and R2 may form a 5- or 6-membered ring; R3-5 = H, halo, alkyl, CO2H, alkoxycarbonyl,

aryloxycarbonyl, amino, carbamoyl, sulfamoyl, NO2, CN, OH, alkoxy, SH, aryl, alkenyl; X1 = COR8, CONR8R9, CO2R8, SO2R8, SOR8, SO2NR8R9; R8,9 = H, (cyclo)alkyl, aryl, heterocyclyl, alkenyl; m = 0-2; Y1 = CN, CONR8R9, CO2R8, SO2R8, SO2R8, SO2NR8R9; X2, Y2 = COR8R9, CO2R8, SO2R8, SOR8, SO2NR8R9].

IT 144806-78-0 144807-06-7 144807-09-0

144807-25-0

RL: USES (Uses)

(bleachable dye, photog. material containing)

RN 144806-78-0 CAPLUS

CN Benzenepropanoic acid, 4-carboxy- α -[[4-(dimethylamino)-2-methoxyphenyl]methylene]- β -oxo-, α -ethyl ester (9CI) (CA INDEX NAME)

RN 144807-06-7 CAPLUS

CN 2-Propenamide, 2-cyano-3-[4-(dimethylamino)-2-methoxyphenyl]-N-[4-[(propylsulfonyl)amino]phenyl]- (9CI) (CA INDEX NAME)

RN 144807-09-0 CAPLUS

CN 2-Propenamide, 3-[2-amino-4-(dimethylamino)phenyl]-N-[4-[(butylsulfonyl)amino]phenyl]-2-cyano- (9CI) (CA INDEX NAME)

RN 144807-25-0 CAPLUS

CN 2-Propenoic acid, 3-[2-methoxy-4-[methyl[2-[(propylsulfonyl)amino]ethyl]amino]phenyl]-2-(phenylsulfonyl)-, ethyl ester (9CI) (CA INDEX NAME)

IT 144807-45-4P

RL: SPN (Synthetic preparation); PREP (Preparation)

(preparation and use of, as bleachable dye, photog. material containing)

RN144807-45-4 CAPLUS

Propanedioic acid, [[4-(dimethylamino)-2-[(methylsulfonyl)amino]phenyl]met CN hylene]-, dibutyl ester (9CI) (CA INDEX NAME)

ANSWER 9 OF 22 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1992:407953 CAPLUS

DOCUMENT NUMBER:

117:7953

TITLE:

Preparation of 4,7-dihydrofuro[3,4-d]pyrimidin-5(1H)-

one derivatives

INVENTOR(S):

Rovnyak, George C.; Kimball, Spencer D.

PATENT ASSIGNEE(S):

E. R. Squibb and Sons, Inc., USA Brit. UK Pat. Appl., 28 pp.

CODEN: BAXXDU

DOCUMENT TYPE:

Patent

LANGUAGE:

SOURCE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION: .

PATENT NO.	KIND	DATE	APPLICATION NO.		DATE
GB 2247236	A1	19920226	GB 1991-17865		19910819 <
GB 2247236	B2	19940105			
US 5103006	Α	19920407	US 1990-570664		19900821 <
PRIORITY APPLN. INFO.:			US 1990-570664	Α	19900821
OTHER SOURCE(S):	MARPAT	117:7953			
GI		•			

Title compds. I [X = O, S; R1 = alkyl, alkenyl, alkynyl,AB (alkyl)cycloalkyl, -aryl, -heterocyclyl, hydroxyalkyl, alkoxyalkyl, mercaptoalkyl, (substituted) amino, heterocyclyl, etc.; R2 = aryl, heterocyclyl] and salts thereof, useful as cardiovascular agents (no data), are prepared Et 4-(acetyloxy)-2-[[2-(methylthio)-3-nitrophenyl]methylene]-3-oxobutanoate (preparation given), 2-methyl-2-thiopseudourea sulfate and AcONa in DMF were heated for 6 h to give an Et (hydroxymethyl)pyrimidinecarboxylate derivative which in MeOH, DMSO and NaOH was stirred at room temperature for 1.5 h to give I [R1 = Me, X = S,

R2

= 2,3-(MeS)(O2N)C6H3]; this was converted to its mono-HCl salt.

IT 141776-01-4P

RL: SPN (Synthetic preparation); PREP (Preparation)

(preparation of, as intermediate in preparation of cardiovascular agents)

RN 141776-01-4 CAPLUS

Butanoic acid, 4-(acetyloxy)-2-[[2-(methylthio)-3-nitrophenyl]methylene]-3-CN oxo-, ethyl ester (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} O \\ \parallel \\ \text{CH} \end{array} \begin{array}{c} C \\ \text{C} \\ \text{C} \end{array} \begin{array}{c} C \\ \text{C} \end{array} \begin{array}{c} C \\ \text{C} \\ \text{C} \end{array} \begin{array}{c}$$

ANSWER 10 OF 22 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1992:106096 CAPLUS

DOCUMENT NUMBER:

116:106096

TITLE:

SOURCE:

GI

Preparation of phenylpyridine derivatives for

treatment of brain and heart ischemia

INVENTOR(S):

Takasugi, Hisashi; Kuno, Atsushi; Sakai, Hiroyoshi

PATENT ASSIGNEE(S):

Fujisawa Pharmaceutical Co., Ltd., Japan Jpn. Kokai Tokkyo Koho, 15 pp.

CODEN: JKXXAF

DOCUMENT TYPE:

Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 03223253 PRIORITY APPLN. INFO.:	A2	19911002	JP 1990-17579 JP 1990-17579	19900126 < 19900126
OTHER SOURCE(S):	MARPAT	116:106096		

Phenylpyridine derivs. [I; R1 = CO2H, alkyl, cyano, alkylsulfonyl, acyl, etc.; R2 = cyano, NO2, halo, (alkyl- or alkoxy-substituted) aryl, heterocyclyl; R3 = (esterified) CO2H, (substituted) carbamoyl, heterocyclylcarbonyl; R4 = alkyl] are prepared BF3-Et2O was added dropwise to a solution of 5 g Et 2-benzoyl-3-(3-nitrophenyl)acrylate in CH2Cl2 at room temperature, followed by a solution of 6.6 g 3-amino-N-(2-morpholinoethyl)crotonamide in CH2Cl2, the mixture was refluxed, the reaction mixture adjusted to pH 9, washed, dried, filtered to give dihydropyridine II, which was refluxed with MnO2 to 1.1 g pyridine derivative III. Also prepared were 33 addnl. I, which restored ATP content by 71.8-93.2% in ischemic guinea pigs at 1 + 10-5 g/mL.

IT 138994-19-1P 138994-20-4P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation and reaction of, in preparation of antiischemic compds.)

RN 138994-19-1 CAPLUS

CN Benzenepropanoic acid, α -[(2-methoxy-5-nitrophenyl)methylene]- β -oxo-, ethyl ester (9CI) (CA INDEX NAME)

RN 138994-20-4 CAPLUS

CN Benzenepropanoic acid, α -[(2-methoxy-3-nitrophenyl)methylene]- β -oxo-, ethyl ester (9CI) (CA INDEX NAME)

ANSWER 11 OF 22 CAPLUS COPYRIGHT 2006 ACS on STN

1991:428892 CAPLUS ACCESSION NUMBER:

115:28892 DOCUMENT NUMBER:

Preparation of phenylalkan(en)oic acids as leukotriene TITLE:

B4 antagonists.

Konno, Mitoshi; Nakae, Takahiko; Hamanaka, Nobuyuki Ono Pharmaceutical Co., Ltd., Japan INVENTOR(S):

PATENT ASSIGNEE(S):

SOURCE: Eur. Pat. Appl., 205 pp.

CODEN: EPXXDW

Patent DOCUMENT TYPE: English LANGUAGE:

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PA	TENT NO.			KINI		DATE	API	PLICATION NO.			DATE	
EP	405116 405116 405116	-		A2 A3 B1		19910102 19920415 19951206	EP	1990-109294			19900516	<- -
	R: AT,	BE,	CH,		DK,	ES, FR,		R, IT, LI, LU,	NL,	SE		÷
	2019335 2019335			AA C		19901227 20000801	CA	1990-2019335			19900507	<
	03261752			A2		19911121	JР	1990-123146			19900515	<
	07039369			В4		19950501						
	619296			A 1		19941012	EP	1994-108324			19900516	<
EP	619296		~~~	B1		19970312	an a		NIT	a r		
ED	R: AT, 652208	BE,	CH,	DE, A1		ES, FR, 19950510		R, IT, LI, LU, 1994-118144		SE	19900516	/
	652208			B1		19980114	111	1994 110144			19900010	`
		BE,	CH,				GB, GI	R, IT, LI, LU,	NL,	SE	E	
AT	131154			E		19951215		1990-109294			19900516	
ES	2083396			TЗ		19960416	ES	1990-109294			19900516	<
AT	150006			E		19970315	AT	1994-108324			19900516	<
ES	2102097			ТЗ		19970716	ES	1994-108324			19900516	<
AT	162181			E		19980115	AT	1994-118144			19900516	<
ES	2114117			Т3		19980516	ES	1994-118144			19900516	<
US	5086065			Α		19920204	US	1990-524521			19900517	<
KR	143404			В1		19980715	KR	1990-7107			19900518	<
US	5155104			Α		19921013	US	1991-760043			19910913	<
US	5256686			Α		19931026	US	1992-921342			19920729	<
JP	06072947			A2		19940315	JP	1993-131187			19930507	<
JP	08019040			В4		19960228						
US	5457122			Α		19951010	US	1993-90456			19930713	<
US	5795914			Α		19980818	US	1995-462815			19950605	
បន	6001877			Α		19991214	US	1998-81549			19980520	<
PRIORIT	Y APPLN.	INFO	.:				JP	1989-164213		Α	19890627	
							JP	1989-310545		Α	19891201	
							JP	1990-1799		A	19900109	
							EP	1990-109294		A3	19900516	
							US	1990-524521		A3	19900517	
							US	1991-760043		АЗ	19910913	
							US	1992-921342		АЗ	19920729	

OTHER SOURCE(S):

MARPAT 115:28892

GI For diagram(s), see printed CA Issue.

AB Title compds. I (A = NHCO, O, NHSO2, CO, CH2, CHOH; W = C1-13 alkylene, phenylene, C6H4CH2; R1 = H, C1-4 alkyl, HO2C, (unsatd.) 4-7-membered N-heterocyclyl, carbamoyl, HOCH2; AWR1 = Q1, Q2, Q3, etc.; Y = CH2CH2, CH:CH; D = hydroxyalkenylene, etc.), are prepared tert-Bu 3-[1-[6-(4-methoxyphenyl)hex-5(E)-enyl]oxy-4-(4-carboxybutanamido)benzen-2-yl]propionate (preparation starting from 2-hydroxy-5-nitrobenzaldehyde given) in THF/Et3N was treated with ClCO2Et at -10° and then with Me2NH to give the dimethylamide derivative which was hydrolyzed in HCO2H to give the title acid-amide E-II. II inhibited binding of 3H-LTB4 to human polymorphonuclear leukocyte LTB4 receptors with IC50 = 0.045 μM. A tablet formulation containing 3-[1-[6-(4-methoxyphenyl)hex-5(E)-enyl]oxy-3-(4-carboxybutyl)oxybenzen-2-yl]propionic acid is given.

IT 134577-68-7P 134577-76-7P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation and reaction of, in preparation of LTB4 antagonists)

RN 134577-68-7 CAPLUS

CN 2-Propenoic acid, 3-(2-hydroxy-5-nitrophenyl)-, 1,1-dimethylethyl ester, (E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 134577-76-7 CAPLUS

CN Pentanoic acid, 5-[[3-[3-(1,1-dimethylethoxy)-3-oxo-1-propenyl]-4-hydroxyphenyl]amino]-5-oxo-, methyl ester, (E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

IT 134578-32-8P

RL: SPN (Synthetic preparation); PREP (Preparation)
 (preparation of, as LTB4 antagonist)

RN 134578-32-8 CAPLUS

CN Pentanoic acid, 5-[[3-.(2-carboxyethenyl)-4-[[6-(4-methoxyphenyl)-5-

hexenyl]oxy]phenyl]amino]-5-oxo-, (E,E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

L5 ANSWER 12 OF 22 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1987:515373 CAPLUS

DOCUMENT NUMBER:

107:115373

TITLE:

Pesticidal 1-(4-aryloxyphenyl)-3-benzoylureas; processes for their preparation, and pesticidal

compositions and methods employing them

INVENTOR(S):

Caruso, Andrew James

PATENT ASSIGNEE(S):

Union Carbide Corp., USA Eur. Pat. Appl., 62 pp.

SOURCE:

CODEN: EPXXDW

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PA'	PATENT NO.						KIND DATE		APPLICATION NO.					
						•								
EP	22084	40			A2		1987	0506	EP	1986-3	07457		19860929	<
EP	22084	40			A3		1988	0323						
	R:	AT,	BE,	CH,	DE,	FR,	GB,	IT,	LI, L	J, NL,	SE			
JP	6211	1961			A2		1987	0522	JP	1986-2	28521		19860929	<
ZA	86074	420			Α		1987	0527	ZA	1986-7	420		19860929	<
AU	86632	260			A1		1987	0402	AU	1986-6	3260		19860930	<
BR	8604	732			Α		1987	0630	BR	1986-4	732		19860930	<
PRIORIT	Y APP	LN.	INFO	:					US	1985-7	81382	A	19850930	
					•				US	1986-8	95364	Α	19860811	

GI

$$X^{3}$$

$$X^{1}$$

$$X^{1}$$

$$X^{2}$$

$$X^{2}$$

$$X^{2}$$

$$X^{2}$$

$$X^{2}$$

$$X^{3}$$

$$X^{2}$$

$$X^{3}$$

$$X^{4}$$

$$X^{5}$$

$$X^{2}$$

$$X^{3}$$

$$X^{2}$$

$$X^{3}$$

$$X^{4}$$

$$X^{5}$$

$$X^{2}$$

$$X^{3}$$

$$X^{4}$$

$$X^{5}$$

$$X^{2}$$

$$X^{3}$$

$$X^{4}$$

$$X^{5}$$

$$X^{7}$$

$$X^{7$$

$$H_2N$$
 O
 $C1$
 $C1$
 $C1$
 $C1$
 $C1$

The title compds. [I; X1 = halo; X2, X3 = H, halo; Y1 = halo, alkyl, alkoxy, NO2, cyano; m = 0-2; R1 = CHO, CO2H or ester, hydroxyalkyl, alkoxyalkyl, acyloxyalkyl, alkenyl, alkanoyl, (a)cyclic acetal, dithioacetal, hemithioacetal; m = 2 and R1 is not at 2- or 6-position when R1 = CO2H or ester; Z = (un)substituted (un)saturated mono- or bicyclic fused ring system (latter has 1 benzene ring and one carbo- or heterocyclic 5- or 6-membered ring containing a CO group and/or 1 or 2 O or S atoms] are prepared as pesticides. Neat 2,6-F2C6H3CONCO (23.63 mmol) was added to a solution of phenoxyaniline derivative II (23.63 mmol) in PhMe. The mildly exothermic reaction precipitated 90% (phenoxyphenyl)benzoylurea III, which was 71-100% lethal against Spodoptera eridania at 100 ppm (spray) on bean leaves in laboratory expts.

III

IT 110123-43-8P
RL: AGR (Agricultural use); BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); USES (Uses) (preparation of, as pesticide)

Ι

RN 110123-43-8 CAPLUS

CN 2-Propenoic acid, 3-[3-chloro-5-[[[(2,6-difluorobenzoyl)amino]carbonyl]amino]-2-(2,4-dimethylphenoxy)-4-methylphenyl]-, methyl ester (9CI) (CA INDEX NAME)

L5 ANSWER 13 OF 22 CAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 1981:603820 CAPLUS

DOCUMENT NUMBER:

95:203820

TITLE:

Addition of heterocyclic CH acids to the carbon-nitrogen double bond of azomethines

AUTHOR(S):

Pavlenko, N. I.; Marshtupa, V. P.; Klyuev, N. A.;

Baskunov, B. P.

CORPORATE SOURCE:

SOURCE:

Donetsk. Gos. Univ., Donetsk, 340055, USSR Khimiya Geterotsiklicheskikh Soedinenii (1981

), (8), 1088-93

CODEN: KGSSAQ; ISSN: 0453-8234

DOCUMENT TYPE:

LANGUAGE:

Journal Russian

GI

Ι

III

IV

Aminomethylation of 1-phenyl-3-methyl-5-pyrazolone by RC6H4CH:NR1 (R = H, 3-NO2, 3-OH, 4-MeO, 2-MeO, 4-Me, 2-HO, 4-Cl; R1 = 4-IC6H4, 4-BrC6H4, 3-BrC6H4, Ph, Me, 7-quinolyl) gave 10-70% addition products I. Treatment of I (R = H, R1 = 4-BrC6H4; R = 4-MeO, R1 = Ph) with acid gave II in 40 and 53% yield, resp. Indolones III (R = 2-OH, 4-OH, 4-Me, 4-MeO, 4-NO2, 4-F, 4-Cl, H, R1 = Et, Ph, 4-O2NC6H4, 4-ClC6H4, 4-BrC6H4, 4-IC6H4) and thiazolidines IV (R = 3-NO2, 4-OH, 4-Me2N, 4-Br, 4-F, 4-NO2, H; R1 = Ph, 3-O2NC6H4, PhOC6H4, Me) were prepared similarly in 31-92% yield. Acid treatment of III gave the corresponding benzylideneindolones. Treatment of IV with OH- gave 15-75% RC6H4CH:C(SH)CO2H.

IT 79787-80-7P

RL: SPN (Synthetic preparation); PREP (Preparation)
 (preparation of)

RN 79787-80-7 CAPLUS

CN 2-Propenoic acid, 3-(2-hydroxy-5-nitrophenyl)-2-mercapto- (9CI) (CA INDEX NAME)

L5 ANSWER 14 OF 22 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1980:568271 CAPLUS

DOCUMENT NUMBER:

93:168271

TITLE:

Hydrazide nucleating agents, methods employing them

and photographic materials containing them

INVENTOR(S):

Sidhu, Jasbir; Simons, Michael John; Baigrie, Brian Devlin; Mijovic, Miroslav Vasa; Southby, David Thomas

PATENT ASSIGNEE(S):

SOURCE:

Kodak Ltd., UK Brit. UK Pat. Appl., 14 pp.

CODEN: BAXXDU

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.		DATE
					10771015
GB 2011391	Α	19790711	GB 1977-52302		19771215 <
GB 2011391	Α	19790711	GB 1978-48701		19781215 <
GB 2011391	B2	19820324			
PRIORITY APPLN. INFO.:			GB 1977-52302	Α	19771215
GI					

AB 3,4-RR1C6H3NHNHCOR2 [I; R = H, R3(Z)nZ1(Z2)m(CH2)x [R3 = group which renders I capable of being adsorbed to the surface of a photog. Ag halide grain; Z, Z2 = divalent aliphatic or aromatic hydrocarbon or heterocyclic moiety; Z1 = NR4CO (R4 = H, alkyl), NR4SO2, O2C, CONR4, SO2NR4, CO2; n, m = 0, 1; x = 1-4]; R1 = R3(O)y (y = 0, 1), R6(CH2)zO (R6 = H, optionally substituted alkyl or aryl, z = 1-4); R2 = H, optionally substituted alkyl or aryl, were prepared Thus, the amide II was prepared (20%) from 5-aminobenzotriazole by stirring it in DMF at room temperature

overnight with p-OCHNHNHC6H4CH2CO2H in the presence of dicyclohexylcarbodiimide. I are useful as photog. nucleating agents. They are adsorbed strongly to Ag halide grains and function at lower pH than previously described. A preferred use of I is in photog. dye image transfer systems both of the peel-apart and integral type.

IT 69447-75-2P

CN

L5 ANSWER 15 OF 22 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1977:552324 CAPLUS

DOCUMENT NUMBER: 87:152324

TITLE: Phosphonium salts and ylides based on chloroacetylurea

AUTHOR(S): Kushnir, V. N.; Shevchuk, M. I.; Dombrovskii, A. V.

CORPORATE SOURCE: Chernovits. Gos. Univ., Chernovtsy, USSR

SOURCE: Zhurnal Obshchei Khimii (1977), 47(8),

1715-21 CODEN: ZOKHA4; ISSN: 0044-460X

DOCUMENT TYPE: Journal

LANGUAGE: Russian

AB Reaction of H2NCONHCOCH2C1 with Ph3P gave 94% H2NCONHCOCH2P+Ph3C1- which on treatment with NH4OH gave 87% H2NCONHCOCH:PPh3 (I). Treating I with RX gave 85-94% H2NCONHCOCHRP+Ph3X- (R = Br, iodo, Me, Me3Si; X = halo) which on dehydrohalogenation gave 67-82% H2NCONHCOCR:PPh3. Treating I with R1CHO gave 77-99% of 16 H2NCONHCOCH:CHR1 (R1 = Ph, substituted phenyl, 2-furyl, 2-quinolyl, etc.) which on bromination gave H2NCONHCOCHBrCHBrR1.

IT 62879-66-7P 62879-67-8P
RL: SPN (Synthetic preparation); PREP (Preparation)

(preparation of)

RN 62879-66-7 CAPLUS

CN 2-Propenamide, N-(aminocarbonyl)-3-(2-hydroxy-3-nitrophenyl)- (9CI) (CA-INDEX NAME)

RN 62879-67-8 CAPLUS

CN 2-Propenamide, N-(aminocarbonyl)-3-(2-hydroxy-5-nitrophenyl)- (9CI) (CA INDEX NAME)

L5 ANSWER 16 OF 22 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1976:91660 CAPLUS

DOCUMENT NUMBER: 84:91660

TITLE: Heterocyclic styryl compounds

INVENTOR(S): Tonegawa, Kakuji; Jono, Shuichi; Fujino, Tomizo PATENT ASSIGNEE(S): Osaka Seika Chemical Industries, Ltd., Japan

SOURCE: Jpn. Tokkyo Koho, 8 pp.

CODEN: JAXXAD

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE	
	JP 50022051	B4	19750728	JP 1966-6348	19660202 <	
PRIO	RITY APPLN. INFO.:			JP 1966-6348	19660202	
GI	For diagram(s), see	printe	d CA Issue.	·		
AB	Styryl fluorescent	whiteni:	ng agents I	(R = H, SO3Na; R1 = H, I	Me; $R2 = H$,	
	Cl, Me or (R2R3) =	benzo; .	A is an opti	onally substituted benze	ene,	
	naphthalene, or heterocyclic ring) are prepared by triazolizing					
	the appropriate ami	no azo	coupling pro-	duct. For example,		

2-(p-aminostyryl)-5-methylbenzoxazole [6661-12-7] was diazotized and coupled with 4,1-H2NC10H6SO3Na [130-13-2] and the product triazolized with NaOCl in aqueous pyridine to give I (R = R1 = R3 = H, R2 = Me, A = 4-sulfo-1,2-naphtho) [58307-08-7], fluorescence λ max 422 m μ . The following I were similarly prepared (R-R3, A, and fluorescence max in m μ given): H, H, H, H, 4-sulfo-1,2-naphtho, 420; H, H, Me, H, 6-sulfo-1,2-naphtho, 440; H, H, Me, H, 7-sulfo-1,2-naphtho, 416; H, H, Me,

6-sulfo-1,2-naphtho, 440; H, H, Me, H, 7-sulfo-1,2-naphtho, 416; H, H, Me, H, 5-sulfo-1,2-naphtho, 449; H, H, Cl, H, 4-sulfo-1,2-naphtho, 421; H, Me, Me, H, 6,8-disulfo-1,2-naphtho, 445; 3-SO3Na, H, H, H, 1,2-naphtho, 429; and 9 others.

IT 58307-05-4

RL: RCT (Reactant); RACT (Reactant or reagent)
 (reaction of, with aminocresol)

RN 58307-05-4 CAPLUS

CN 2-Propenoic acid, 3-(4-nitro-2-sulfophenyl)- (9CI) (CA INDEX NAME)

L5 ANSWER 17 OF 22 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1974:84703 CAPLUS

DOCUMENT NUMBER: 80:84703

TITLE: Yellow coumarin dyes

INVENTOR(S):
Sato, Katsunobu

PATENT ASSIGNEE(S): Sumitomo Chemical Co., Ltd. SOURCE: Jpn. Kokai Tokkyo Koho, 5 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 48080122	A2	19731026	JP 1972-11985	19720201 <
JP 51042611	B4	19761117		

PRIORITY APPLN. INFO.: JP 1972-11985 A 19720201

Coumarin dyes (I, R1, R2 = H, alkyl, or cycloalkyl, or R1, R2, and N form a heterocyclic group; X = S, NH, or NR3, R3 = alkyl, aryl, or aralkyl; A = benzene or naphthalene ring with or without substituents except CO2H and SO3H) are prepared through condensation reactions. The dyes are useful for dyeing acetate, polyester, or polyamide fibers in fluorescent yellow shades with good fastness. Thus, NCCH2CONH2 was treated with 4,2-(Et2N)(HO)C6H3CHO in MeOH containing piperidine at room temperature

to give 4,2-(Et2N) (HO)C6H3CH:C(CN)CONH2 which was treated with o-(H2N) 2C6H4 in DMF at 100-10.deg. to give a yellow dye (I, R1 = R2= Et, X = NH, A = benzene ring) [27425-55-4]. Similarly prepared were 2 other I.

TT 42005-48-1P RL: IMF (Industrial manufacture); PREP (Preparation) (preparation of)

42005-48-1 CAPLUS RN

2-Propenamide, 2-cyano-3-[4-(diethylamino)-2-hydroxyphenyl]- (9CI) (CA CN INDEX NAME)

$$\begin{array}{c|c} \text{Et}_2\text{N} & \text{NC} & \text{O} \\ & | & | \\ \text{CH} \hline & \text{C} - \text{C} - \text{NH}_2 \end{array}$$

ANSWER 18 OF 22 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1972:128826 CAPLUS

DOCUMENT NUMBER:

76:128826

TITLE:

Oxazolylacetic acid derivatives and oxazolylcoumarins

for dyeing organic fibers

INVENTOR(S):

Harnisch, Horst

PATENT ASSIGNEE(S):

Farbenfabriken Bayer A.-G.

SOURCE:

Ger. Offen., 80 pp.

CODEN: GWXXBX

DOCUMENT TYPE:

Patent

LANGUAGE:

German

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PA	TENT NO.	KIND	DATE	APP	LICATION NO.	DATE	
DE	2030507	A	19720105	DE	1970-2030507	19700620	<
DE	2030507	B2	19740919				
DE	2030507	C3	19750522				
CH	717157	A4	19760630	CH	1971-7157	19710513	<
CH	587833	Α	19770513	CH	1973-16185	19710513	<
CH	585250	Α	19770228	CH	1973-16186	19710613	<
BE	768722	A1	19711103	BE	1971-104800	19710618	<
NĻ	7108436	Α	19711222	NL	1971-8436	19710618	<
FR	2099247	A5	19720310	FR	1971-22352	19710618	<
GB	1329042	A	19730905	GB	1971-28704	19710618	<
GB	1329043	Α	19730905	GB	1972-38453	19710618	<
AT	310707	В	19731010	AΤ	1971-5278	19710618	<
AΤ	310743	В	19731010	AΤ	1972-6152	19710618	<

JP 50023028	В4	19750805	JP 1971-43359		19710618 <
US 3985763	Α	19761012	US 1973-369124		19730612 <
JP 50069380	A2	19750610	JP 1974-99075		19740830 <
JP 51006266	В4	19760226			
JP 51000526	A2	19760106	JP 1974-99076		19740830 <
JP 51042125	B4	19761113			
PRIORITY APPLN. INFO.:			DE 1970-2030507	Α	19700620
			DE 1970-2058877	Α	19701130
			US 1971-154652	A1	19710618

Oxazoles [I, A represents benzene, naphthalene, or dibenzofuran ring; R = AB H, alkyl, cyclohexyl, aralkyl, aryl; R1 = H, alkyl, cyclohexyl, aralkyl, aryl, or (RR1N) = heterocyclic ring] were prepared by reaction of o-aminophenols with NCCH2CONRR1 and treated with 4-(dialkylamino)salicylaldehydes to give oxazolylcoumarins (II, R = Me, Et), fluorescent dyes for natural and synthetic fibers. For example, a mixture of o-H2NC6H4OH and NCCH2CONH2 was heated under N 30 min at 140-60.deg., 15 min at 150-60.deg., and 1 hr at 170.deg. to give 2-(2benzoxazolyl)acetamide [34564-12-0]. Similarly, 46 other I were prepared A mixture of NCCH2CO2Et and MeO(CH2)3NH2 was heated 30 min at 60.deg., 3,4-H2N(HO)C6H3Me added, and the mixture heated 6 hr at 180.deg. to give N-(3-methoxypropyl)-5-methyl-2-benzoxazoleacetamide which (without isolation) was refluxed 20 hr with 4,2-Et2N(HO)C6H3CHO and iso-PrOH in the presence of piperidine to give 7-(diethylamino)-3-(5-methyl-2benzoxazolyl)coumarin [34564-13-1], dyeing nylon-6 fabric a fast, brilliant greenish yellow shade. Similarly, 13 other II were prepared IT 35773-52-5P

RL: IMF (Industrial manufacture); PREP (Preparation)
 (preparation of)

RN 35773-52-5 CAPLUS

CN 2-Benzoxazoleacetamide, α-[[4-(diethylamino)-2ethoxyphenyl]methylene]-N,5-dimethyl- (9CI) (CA INDEX NAME)

L5 ANSWER 19 OF 22 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1944:41982 CAPLUS

DOCUMENT NUMBER: 38:41982

ORIGINAL REFERENCE NO.: 38:6288b-i,6289a-c

TITLE: .Nitrogen heterocycles. LI. A new linear

benzodipicolone, 2,6-dimethyl-1,5-anthrazoline

(2,7-dimethylpyrido[2,3-g]quinoline)

AUTHOR(S): Ruggli, Paul; Brandt, Fritz

SOURCE: Helvetica Chimica Acta (1944), 27, 274-91

CODEN: HCACAV; ISSN: 0018-019X

DOCUMENT TYPE: Journal LANGUAGE: Unavailable

OTHER SOURCE(S): CASREACT 38:41982

AB cf. C. A. 37, 1714.8. The successful use of 4,6-diaminoisophthalaldehyde for the previous synthesis of 1,8-anthrazoline derivs. (C. A. 32, 562.7)

for the previous synthesis of 1,8-anthrazoline derivs. (C. A. 32, 562.7) suggested the use of the corresponding 2,5-diaminoterephthalaldehyde (I) as a starting material for the preparation of derivs. of a new linear benzodipicoline, 2,7-dimethylpyrido[2,3-g]quinoline (II). Attempts to prepare I through 2,5-dichloroterephthalaldehyde (III) from

2,5-dichloro-p-xylene (IV) are briefly described though the yields and purity of the products left so much to be desired that a more successful approach was made through the corresponding 2,5-dibromoterephthalaldehyde (V). The chlorination of 50 g. p-xylene in the presence of 5 g. Fe powder in the dark at 12-15° in 3 hrs. and crystallization of the product from MeOH gave 43 g. (50%) of IV, m. 70-1°. Chlorination of the side-chain by passing dry Cl into 20 g. IV in 12 g. C6H2Cl4 at 120-30° with illumination gave 16.8 g. of 1,4-bis(dichloromethyl)-2,5-dichlorobenzene (VI), m. 72.5-4.0°, saponified by heating at 150-70° with concentrated H2SO4 for 20 min. The crude product, m. 144°, was purified through the dianil, C20H14Cl2N2, m. 213-140°, saponified by refluxing with 10% HCl and recrystd. from PhNO2 to give yellow needles of III, m. 157-8°. Other chlorination products including 2,3,5,6-tetrachloro-p-xylene, m. 216.5-17.0°; 1,4-bis (chloromethyl)-2,3,5,6-tetrachlorobenzene, m. 174.5-5.0° (dianil, C20H16Cl4N2, m. 170°); 1,4-bis(trichloromethyl)-2,5-dichlorobenzene, m. 193°. Bromination of 25 g. IV at 180 with 92 q. Br for 3.5 hrs. and crystallization of the product from CHCl3 yielded 40 g.

of

1,4-bis(dibromomethyl)-2,5-dichlorobenzene, m. 127.5-8.0°. The bromination of 20 g. of p-xylene at $10-15^{\circ}$ in the presence of a trace of iodine with 21.1 cc. Br and recrystn. of the crude product from alc. gave 44 g. of 2,5-dibromo-p-xylene (VII), m. 73.5-4.0°. Bromination of the side chain by adding in 5 hrs. 42.5 cc. Br to 50 g. VII at 120° and recrystn. of the crude product from 1100 cc. of boiling AcOEt yielded 78-81 g. (71-4%) of light yellow needles of $\alpha, \alpha, \alpha', \alpha', 2, 5$ -hexabromo-p-xylene (VIII), m. 160-2. A mixture of 50 q. VIII and 250 cc. of H2SO4.H2O was heated at 130-40° and 25 mm. for 1 hr. The cooled solution was diluted with 1 kg. of ice and the crude product (26 g., m. 180-5°) was recrystd. from 250 cc. AcOH, producing 21.1 g. (84%) of V, m. 189-190.5°; dianil, m. 234.5-5.0°; tetraacetamide, m. above 305°. A mixture of 10 g. V with 1 g. Cu powder, 1 g. CuBr, 1 g. K2CO3, 18 g. of p-MeC6H4SO2NH2 and 40 cc. PhNO2 was heated at 140° and treated with 14 g. K2CO3 in 2 hrs. at 150-5°. After 3 hrs. at 160° the reaction mass was worked up and the crude product was recrystd. from AcOH and PhNO2, yielding 52-4% of 2,5-di-p-tolylsulfonamidoterephthalaldehyde (IX), C22H2ON2O6S2, m. 241-3° (decomposition); dianil, m. 297° (decomposition). Condensation of 5 g. IX with 25 cc. AcCH2CO2Et at 70 in the presence of 12 drops of piperidine and crystallization from alc. gave 90% of

di-Et

2,5-bis(p-tolylsulfonamido)terephthalylidenediacetoacetate (X), ${\tt C34H36N2O10S2,\ m.\ 216-17}^{\circ}$ (decomposition). Treatment of 1.5 g. X with 5 cc. concentrated ${\tt H2SO4}$ at ${\tt 27-32}^{\circ}$ (not over 40) gave a one-sided ring-closure with the formation of Et 2-methyl-3-carbeth oxy-6-amino-7-quinoline (methyleneacetoacetate) (XI), m. 219-20°; picrate, m. 215-20° (decomposition). Treatment of 5 g. X with 20 cc. H2SO4 for 1 hr. below 95° saponified the ester group and gave 1.9 g. of 2-methyl-3-carboxy-6-amino-7-quinoline(methyleneacetoacetic acid) (XII) which on further treatment with concentrated H2SO4 at 98-100° underwent further ring closure to 2,7-dimethylpyrido[2,3-g]quinoline-3,8dicarboxylic acid (XIII), m. 320° (decomposition), also similarly prepared from X and XI. Decarboxylation of 1 g. XIII by adding it portionwise in 5 min. to 12 cc. quinoline at 215° containing 0.2 g. Cu powder and 0.2 g. CuCrO2, followed by removal of the quinoline with steam distillation and recrystn. of the crude product from alc., gave 0.2 g. (30%) of needles of II, C14H12N2, m. 238-9° (decomposition); picrate, m. 263° (decomposition); dibenzylidene derivative, m. 267°; bis(pdimethylaminobenzylidene) derivative, m. above 340°. Treatment of 0.5 g. IX with 5 cc. PhCOMe at 190-7° for 1.5 hrs. and recrystn. of the product from alc. and PhNO2 produced greenish yellow leaflets of 2,7-diphenylpyrido[2,3-g]quinoline, m. 284-5°. From 100 g.

p-xylene, the main products were 123 g. aldehyde (V), 100 g. sulfonamide (IX), 105 g. condensation product (X), 25 g. dicarboxylic acid (XIII) and, finally, 5 g. II.

IT 857619-45-5, Acetoacetic acid, α,α' -[2,5-bis(p-tolylsulfonamido)terephthalylidene]bis-, diethyl ester (preparation of)

RN 857619-45-5 CAPLUS

CN Acetoacetic acid, $\alpha, \alpha'-[2,5-bis(p-tolylsulfonamido) terephthalylidene]bis-, diethyl ester (4CI) (CA INDEX NAME)$

L5 ANSWER 20 OF 22 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1940:12844 CAPLUS

DOCUMENT NUMBER: 34:12844

ORIGINAL REFERENCE NO.: 34:1986a-i,1987a

TITLE: Nitrogen heterocycles. XLVI.

4,6-Diaminoisophthalaldehyde. 3

AUTHOR(S): Ruggli, Paul; Frey, Hugo

SOURCE: Helvetica Chimica Acta (1939), 22, 1413-27

CODEN: HCACAV; ISSN: 0018-019X

DOCUMENT TYPE: Journal LANGUAGE: Unavailable

The 3,6-dicarboxylic ester produced by the addition of 2 mols. AcCH2CO2Et to 4,6-diaminoisophthalaldehyde (I) was saponified to the free acid which was decarboxylated by heating with Cu in quinoline at 160-230° for 20 min. The resulting 2,7-dimethylbenzodipyridine (II) was converted into the hexa-Br derivative which was transformed by heating with oleum to the crude benzodipyridine-2,7-dicarboxylic acid (III). A mixture of 0.25 g. III, 2 cc. of 10% NH4OH and 2 cc. alc. was triturated, diluted with 20 cc. H2O and heated. The NH3-free product was diluted with 10 cc. H2O and boiled with 0.5 g. AgNO3 in 10 cc. H2O. The crude Ag salt (0.45 g.) was boiled with 70 cc. MeOH and 0.4 g. MeI for 1 h., filtered and concentrated to 20 cc., yielding 0.2 g. (70%) of yellow needles of di-Me benzodipyridine-2,7dicarboxylate, C16H12N2O4, m. 272° (with darkening). Decarboxylation of III gave benzodipyridine (IV); perchlorate, m. 268° (explosive on rapid heating); MeI derivative, m. above 200° (decomposition). Reduction of 0.2 g. IV in 5 cc. of boiling AmOH with 0.35 g. Na

and recrystn. from alc. gave octahydrobenzodipyridine, m. 111.5°, identified through the di-NO and di-Ac derivs., m. 179° (decomposition) and 143°, resp. Reduction of II with Na in AmOH gave as main product a resin which was converted into a colorless crystalline octahydro-2,7-dimethylbenzodipyridine diperchlorate, C14H22Cl2N2O8, m. 285-6° (decomposition). The resinous free base yielded 2 isomeric di-NO derivs., m.

164.5 and 151.5-2.0°, resp. Condensation of 0.2 g. II with 0.5 g. of p-Me2NC6H4CHO at $170-5^{\circ}$ in the presence of 10 drops of piperidine produced 0.45 g. of orange-red 2,7-bis(pdimethylaminostyryl)benzodipyridine, C32H30N4, m. about 340° (with darkening), dissolving in HCl to give violet, blue, green and yellow solns. with increasing acid concns. Condensation of II with o-C6H4(CO2Et)2 by heating in the presence of Na for 14 h. at 100° gave a scarlet crystalline powder which on sulfonation dyed wool and silk bluish red in an acid bath. A unilateral condensation of 0.6 g. I with 6 cc. AcCH2CO2Et occurred on heating in the presence of 9 drops of piperidine for 30 min. at 170°. The impure 3-acetyl-6-formyl-7aminocarbostyril yielded yellow crystals of a pure Ac derivative, C14H12N2O4, m. 320-40° (decomposition). Treatment of 1 g. I in 100 cc. alc. at 30° with 14 g. of dry OHCCHNaCO2Et, boiling for 1 h. after standing for 3 days, filtering off the brown amorphous precipitate (V), adding 1 cc. H2O and standing for 8 days gave a Na salt which was dissolved in 50 cc. H2O, acidified with 10% HCl and recrystd. from dioxane, yielding di-Et 2,6-diaminoisophthalaldiformylacetate, C18H2ON2O6, m. 250°

(decomposition). V was dissolved in H2O, filtered and precipitated with dilute HCl.

The amorphous product (0.06 g.) was decarboxylated by heating in vacuo with 0.3 g. BaO and 0.5 g. Cu at 150° to yield a bright yellow sublimate of IV. Condensation of I with excess cyclohexanone in the presence of piperidine produced 2,3,6,7-bis (tetramethylene)-benzodipyridine, C20H2ON2, m. 250-1° (with darkening); dipicrate, m. 195° (decomposition). A mixture of 8 g. I in 150 cc. alc., 24 cc. PhCH2CN and 12.5 cc. of 30% NaOH was heated for 30 min. on the steam bath. Working up and purification through the di-HCl salt gave a free base (VI), C24H18N4, m. 301°; tetra-Ac derivative, C32H26N4O4, m. 238.5-9.5° (decomposition). Saponification of VI with HCl produced a carboxyl derivative, C24H18N2O3, which gave a Na salt and a mono-Ac derivative, m. 365°. Condensation of 4 g. of 4,6-dinitroisophthalaldehyde with 8.4 q. of dry PhCH(Na)CO2H by heating with 34 cc. Ac2O and 1.2 g. ZnCl2 for 40 h. at 80° gave a powdery dicarboxylic acid which was esterified through the Ag salt to di-Me 4,6-dinitroisophthalalbis(phenylac etate), C26H2ON2O8, m. 152.5-3.5°. Condensation of methazonic acid (VII) with o-H2NC6H4CHO yields 3-nitroquinoline and similarly a cold mixture of VII and I in the presence of a min. of HCl gave 20% of yellow-orange needles of a compound C16H14N6O5, m. 290° (decomposition), of undetd. composition

1T 857578-13-3, m-Benzenediacrylic acid, 4,6-diamino- α , α '-diformyl-, diethyl ester (preparation of)

RN 857578-13-3 CAPLUS

CN m-Benzenediacrylic acid, 4,6-diamino- α , α '-diformyl-, diethyl ester (4CI) (CA INDEX NAME)

L5 ANSWER 21 OF 22 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1939:8734 CAPLUS

DOCUMENT NUMBER: 33:8734

ORIGINAL REFERENCE NO.: 33:1325a-i,1326a

Nitrogen heterocycles. XXXV. 4,6-Dinitro-TITLE:

and diaminoisophthalaldehydes. 2. $lin-Benzodi-\alpha-$

picoline and benzodipyridine

AUTHOR(S): SOURCE:

Ruggli, Paul; Hindermann, Peter; Frey, Hugo Helvetica Chimica Acta (1938), 21, 1066-83

CODEN: HCACAV; ISSN: 0018-019X

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Unavailable

Journal

LANGUAGE: cf. C. A. 32, 3394.4. Dinitroisophthalaldehyde (I) (7 g.) in 40 cc. pyridine was warmed to 60°. CO2 and nitrous fumes developed, the temperature rose to 100° and the reaction ended in 45 min. Recrystn. of the resulting 4.8 g. of brown powder gave yellow leaflets, C25H18N2O6, m. above 300°. Other reactions of I with barbituric acid, indandione and methylphenylpyrazolone are cited. The product (0.5 g.) of the reaction between 7 g. I and CH2N2 (C. A. 31, 4287.9) is now considered to be 4,6-dinitrophenylene-1,3-diethylene oxide; C10H8N2O6, m. 153-4°, converted by HCl in pyridine to the corresponding 4,6-dinitrophenylene-1,3-diethylene chlorohydrin, C10H10Cl2N2O6, m. 150-1°. Boiling 2 g. Et diaminophenylenediacrylate (C. A. 31, 4287.9) with 30 cc. concentrated HCl for 15 min. gave 1.2-1.4 g. of impure 4,6-diaminophenylene-1,3-diacrylic acid HCl salts (II), converted by heating with a 20-fold excess of Ac2O at 120° to the mono-Ac derivative, C14H14N2O5, m. 320° (decomposition). Refluxing with 80 parts Ac20 for 50 min. produced the di-Ac compound, C16H16N2O6, m. 320° (decomposition). The mother liquors of the above saponification yielded yellow matted needles of 7-aminocarbostyril-6-acrylic

acid,

C12H10N2O3, m. above 300°. Heating 0.5 g. II with 25 cc. concentrated HCl in a bomb-tube for 5 h. at 160° gave, by double ring-closure, 2,7-dihydroxybenzodipyridine, C12H8N2O2, charring above 400°. Most condensations run more smoothly with diaminoisophthalaldehyde (III) than with I, on account of the sensitivity of the latter to alkaline condensation agents. Thus, refluxing 0.65 g. III in 50 cc. alc. and 1 g. barbituric acid in 30 cc. H2O for 10 min. produced 1.4 g. of fine, crystalline orange powder, 4,6-diaminoisophthalaldibarbituric acid, C16H12N6O6, charring above 300°. It is remarkable that no further ring-closure between the adjacent CO and NH2 groups takes place as in the condensation of o-H2NC6H4CHO with barbituric acid. In the presence of 10 drops of KOH in MeOH 0.5 g. III condensed with 5 g. of p-MeOC6H4Ac at 150° to give 0.6 g. of 2,7-di(p-methoxyphenyl)benzodipyridine, C26H2ON2O2, m. 268-9°. Condensation of III (2.5 g.) with 10 g. AcCH2Ac in the presence of 15 drops of piperidine in a bomb-tube at 180-90° for 1.5 h. gave 3.5 g. of 2,7-dimethyl-3,6-diacetylbenzodipyridine dihydrate, C18H16N2O2.2H2O, m. 213-15°, converted by heating with Ac2O for 1 h. into an addition compound, C18H16N2O2.Ac2O which, on warming, gave the free base; dioxime, C18H18N4O2, m. 255-7°. III condensed with BzCH2CO2Et by 1-sided ring condensation to 3-benzoyl-6-aldehydo-7aminocarbostyril, C17H12N2O3, m. 278-9° (decomposition); Ac derivative, C19H14N2O4, m. about 320° (decomposition). The ester resulting from the condensation of III with AcCH2CO2Et in the presence of alc. NaOH (C. A. 31, 4287.9) was saponified and decarboxylated by heating 10 g. of the ester with 75 cc. concentrated HCl in a Durobax bomb-tube (70 cm. by 2.2 cm.; capacity, 270 cc.) up to 130° in 1.0-1.5 h. and for 2 h. at 130°. The crude product gave a high-melting polymer, C14H12N2.2H2O, m. 268°, and 2.8 g. of benzodi- α -picoline (IV), C14H12N2, m. 196-7°; dipicrate, m. 220° (decomposition); monoperchlorate, m. 228-30° (decomposition); diperchlorate, m. 318° (decomposition); chromate; MeI compound, sintering at 244°; dibenzal derivative, C28H2ON2, m. 279°; difural derivative, C24H16N2O2, m. 271.5-2.5° (decomposition). Bromination of 4 g. IV in 80 cc. AcOH and 20 g. anhydrous AcONa at 70° with 18.5 g. Br in 40 cc. AcOH with stirring gave 12 g. (90%) of the hexa-Br derivative (V), C14H6Br6N2, m. 190-2° (decomposition), converted by heating with 15% oleum for 50 min.

into the corresponding dicarboxylic acid (VI). A mixture of 0.6 g. VI, 2.5 q. Naturkupfer C, 1.8 g. anhydrous Ba(OH)2 and 1.8 g. BaO was sublimed in vacuo at 230-40° and yielded 45% (1.8 g.) of a yellow crystalline sublimate, m. 159-63°. The crude was dissolved in 15 cc. CHCl3 (distilled over K2CO3), filtered and shaken out with 2 cc. of 10% NaOH and with 4 lots of H2O (3 cc.). After drying over MgSO4, treating with charcoal and evaporating, the residue (0.11 g.) was recrystd. from 8 cc. H2O to give snow-white needles of lin-benzodipyridine (1,8-diazaanthracene), C12H8N2, m. 164.5-5.0°; dipicrate, m. 262° (darkening).

857578-15-5, m-Benzenediacrylic acid, 4,6-diamino-ΙT

(hydrochlorides)

RN 857578-15-5 CAPLUS

m-Benzenediacrylic acid, 4,6-diamino- (4CI) (CA INDEX NAME) CN

857578-17-7, m-Benzenediacrylic acid, 4,6-diacetamido-857578-20-2, m-Benzenediacrylic acid, 4-acetamido-6-amino-(preparation of)

857578-17-7 CAPLUS RN

m-Benzenediacrylic acid, 4,6-diacetamido- (4CI) (CA INDEX NAME) CN

RN 857578-20-2 CAPLUS

m-Benzenediacrylic acid, 4-acetamido-6-amino- (4CI) (CA INDEX NAME) CN

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1937:30573 CAPLUS ACCESSION NUMBER:

31:30573 DOCUMENT NUMBER:

31:4287i,4288a-f ORIGINAL REFERENCE NO.:

Nitrogen heterocycles. XXVIII. TITLE:

4,6-Dinitro-and diaminoisophthalaldehyde. 1

Ruggli, Paul; Hindermann, Peter

AUTHOR(S):

Helvetica Chimica Acta (1937), 20, 272-82 SOURCE:

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4,6-Dinitro-1,3-xylene (100 g.) and 150 g. p-NOC6H4NMe2 were boiled 8 h. in 500 cc. EtOH containing 100 g. anhydrous Na2CO3. Extraction of the crude product

with 1.5 l. H2O and then 3 times with 350 cc. Me2CO left 57% of condensation product (I), 100 g. of which was shaken 24 h. with 620 cc. C6H6 (II) and 620 cc. HNO3 (d. 1.12). After filtering off the p-NH2C6H4NMe2.HNO3, the II layer was separated, and concentrated to 100 cc., when

4,6-dinitroisophthalaldehyde (III) (dianil, m. 164.5-65°; disemicarbazone, m. above 360° (decomposition)) crystallized III condenses with compds. containing an active CH2 group. Thus 1.5 g. III in 10 cc. pyridine (IV) was added to 3 g. barbituric acid in 90 cc. hot H2O. After long standing addition of dilute H2SO4 precipitated 4,6-dinitroisophthalaldibarbituric

acid. CH2N2 (from 23 g. NO(Me)NCO2Et) in 200 cc. ether was poured over 7 g. III and left 15 h. in the ice box. Long fractional crystallization of the precipitate

from EtOH gave 4,6-dinitro-1,3-diacetylbenzene, m. 153-4°. III (20 g.), 100 g. (HO2C)2CH2 and 60 cc. IV were warmed 48 h. at 50-5° and then 2 h. at 100°. Addition of 300 cc. 10% H2SO4 gave 68% of 4,6-dinitrophenylene-1,3-diacrylic acia, m. 216°, after purification through the Et ester (V), m. 116°, and saponification with H2SO4 in dilute AcOH. Reduction of 18 g. V with Rupe's Ni catalyst (VI) gave 14 g. di-Et 4,6-diaminophenylene-1,3-diacrylate, m. 195-6° (di-Ac derivative, m. 244-5°). Reduction of III with VI was unsuccessful. III (16 g.) in 600 cc. EtOH and 360 cc. concentrated NH4OH was dropped with strong stirring during 15 min. into 368 g. FeSO4 in 800 cc. H2O containing a few drops of 10% HCl warmed on the water bath. The Fe precipitate was extracted 15 h. in a Soxhlet

with Me2CO (VII) and the residue after removal of VII, boiled with H2O and filtered. On strong chilling 84% of 4,6-diaminoisophthalaldehyde (VIII), m. 208°, separated; dioxime, m. 219-20°; disemicarbazone, chars above 360°; monophenylhydrazone, m. 275-6° (decomposition); diphenylhydrazone, m. 337° (decomposition); mono-Ac derivative, from VIII and Ac2O in the cold for 3 days, m. 270-2°; di-Ac derivative, prepared hot, m. 280-2°. VIII (0.5 g.) in 5 cc. MeCOPh containing 3-4 drops 10% MeOH-KOH at 100° for 10 min. gave, on precipitation with 50% EtOH, 70% of 2,7-diphenyl-lin-m-benzodipyridine, m. 216-17° (dipicrate, m. 270° (decomposition)). Similar condensation of VIII with AcCH2CO2Et gave di-Et 2,7-dimethylbenzodipyridine-3,6-dicarboxylate, m. 166-7°.

IT 857578-14-4, m-Benzenediacrylic acid, 4,6-diamino-, diethyl ester 857578-16-6, m-Benzenediacrylic acid, 4,6-diacetamido-, diethyl ester

(preparation of)

RN 857578-14-4 CAPLUS

CN m-Benzenediacrylic acid, 4,6-diamino-, diethyl ester (4CI) (CA INDEX NAME)

$$\begin{array}{c} O \\ \parallel \\ E t O - C - C H = C H \\ \\ H_2 N \\ \end{array} \begin{array}{c} O \\ \parallel \\ C H = C + C - O E t \\ \\ N H_2 \end{array}$$

RN 857578-16-6 CAPLUS

CN m-Benzenediacrylic acid, 4,6-diacetamido-, diethyl ester (4CI) (CA INDEX NAME)